PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



(51) International Patent Classification 6:		(1	1) International Publication Number: WO 98/4401
C08F 210/16, 2/34, 4/602, C08J 5/18	A1	(4:	3) International Publication Date: 8 October 1998 (08.10.98
(21) International Application Number: PCT/US (22) International Filing Date: 27 March 1998 ((81) Designated States: AU, BR, CA, CN, JP, KR, MX, SC Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM European patent (AT, BE, CH, DE, DK, ES, FI, FR, GF, GR, IE, IT, LU, MC, NL, PT, SE).
(30) Priority Data: 60/042,310 1 April 1997 (01.04.97)	τ	JS	Published With international search report.
(71) Applicant: EXXON CHEMICAL PATENTS INC. 5200 Bayway Drive, Baytown, TX 77520-5200 (1		S];	
(72) Inventors: LUE, Ching-Tai; 14938 Redwood Be Houston, TX 77062 (US). MERRILL, Natalie, Aurora Street, Houston, TX 77008 (US). MUHLE, E.; 5902 Leafy Aspen Court, Kingwood, TX 773 VAUGHAN, George, A.; 3606 Durhill Street, Hou 77025 (US).	, A.; 60 , Michae 345 (US	01 : el, S).	
(74) Agents: HARRIS, Gerald, L. et al.; Exxon Chemical C P.O. Box 2149, Baytown, TX 77522-2149 (US).	Compan	ıy,	
(54) Title: IMPROVED EASY PROCESSING LINEAR	LOW I	DEN	NSITY POLYETHYLENE
atoms obtainable by a continuous gas phase polymerization using supported	as a met	tallo	O+e (11.71-0.000268xM+2.183x10 ⁻⁹ xM ²)] (I) cene in the substantial absence of an aluminum alkyl based scavenge to 15; a Compositional Distribution Breadth Index (CDBI) as defin
which polymer has a Melt Index (MI) as herein defined or		TI	e value as herein defined of less than 20 %; a Melt Index ratio (MI
which polymer has a Melt Index (MI) as herein defined of herein of at least 70 %, a density of from 0.910 to 0.930 g as herein defined of from 35 to 80; an averaged Modulus a relation bewteen M and the Dart Impact Strength in g/m	g/ml; a l (M) as l	here	in defined of from 20 000 to 60 000 psi (pounds per square inch) a omplying with formula (I).
which polymer has a Melt Index (MI) as herein defined of herein of at least 70 %, a density of from 0.910 to 0.930 g as herein defined of from 35 to 80; an averaged Modulus	g/ml; a l (M) as l	here	sin defined of from 20 000 to 60 000 psi (pounds per square inch) a omplying with formula (I).
which polymer has a Melt Index (MI) as herein defined of herein of at least 70 %, a density of from 0.910 to 0.930 g as herein defined of from 35 to 80; an averaged Modulus	g/ml; a l (M) as l	here	in defined of from 20 000 to 60 000 psi (pounds per square inch) a smplying with formula (I).

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑŪ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
ВJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
СН	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		
]							
İ				_			

WO 98/44011 PCT/US98/06081

IMPROVED EASY PROCESSING LINEAR LOW DENSITY POLYETHYLENE

FIELD OF THE INVENTION

The invention relates to novel polyethylene compositions and particularly to polyethylene having improved combination of shear thinning behavior (to assist in the processing of such polymers in the molten state) and impact strength (to assist the end-use performance). The polymers can be optimally produced in a continuous gas phase processes in which supported catalyst is introduced into a fluidized bed reactor.

BACKGROUND OF THE INVENTION

Polyethylene produced from gas phase processes with a degree of branching to improve melt rheology are described in EP-A-495099; EP-A-452920; EP-A-676421 and EP-A-659773. WO 96/08520 (Exxon Chemical Patents Inc) discusses gas phase polymerization in a low scavenger environment, that is to say no or only a low amount of scavenger in the form of, for example, triethyl aluminum, is used in the course of polymerization.

20

Polyethylene with improved rheology obtained with monocyclopentadienyl compounds are described in WO-A- 93/08221.

EP-A-495099 produces polyethylene (see page 6) using hafnium metallocene compounds having multidentate ligands (i.e. they have two cyclopentadienyl ring systems connected by a bridge). The specifically named hafnium compounds are bridged. The described polymerization is performed in a batch system. The polymerization is performed with unsupported catalyst in a solution phase although page 13 line 12 does mention vapor phase operation. The properties of the resulting polyethylene include a narrow molecular weight distribution and a Melt

15

Flow Rate (MFR expressed in g per 10 minutes at 190 C under a load of 2.16 kg) of from 8-50. The abbreviation MFR is used to indicate Melt Flow Rate or Melt Flow Ratio depending on the source. Reference must be made to the original source in case of doubt to determine the meaning of MFR in a particular case.

EP-A-452920 does exemplify the use of supported catalyst for making polyethylene. The polymerization is in the gas phase using triisobutyl aluminum as a scavenger. The transition metal component includes zirconocenes. Example 9 and others use ethylene-bridged bis(indenyl)zirconium as the transition metal compound. Example 10 uses an Al/Zr ratio of 112. The scavenger helps to avoid the effect of adventitious poisons attached to the experimental equipment or introduced with the various components. The melt tension is said to be improved.

WO-A-95/07942 uses monocyclopentadienyl compounds in a gas phase on a support for producing polyethylene. The activator is not methyl alumoxane but a non-coordinating bulky anion first described in EP-A-277003 and EP-A-277004. Polymerization was performed in a batch reactor. Scavenger was not mentioned.

US-A-5466649 describes in Example 17 preparing polyethylene using a batch gas phase polymerization procedure using dimethylsilyl bis(tetrahydroindenyl) zirconium dichloride on one support and TMA (trimethyl aluminum) supported separately on another support. This was a batch reaction and no detailed indication of the polyethylene properties was given.

WO 96/08520 (Exxon Chemical Patents Inc) describes a continuous commercial gas phase operating process in which scavenger is either not

PCT/US98/06081

present or present in a reduced amount. One embodiment (see page 12, line 28) defines a system essentially free of scavenger, i.e. containing less than 10 ppm of scavenger based on the total weight of the feed gas, which is there referred to as the recycle stream. Alternatively, the low scavenger condition is defined in relation to the metallocene. On page 14, a molar ratio is defined of from 300 to 10. On page 15 it is indicated that the number of olefinic or unsaturated oligomers in the resulting polymer is greatly reduced.

EP-A-676421 (BP) exemplifies a batch type process and a continuous process for producing polyethylene which leads to an improved rheology product through introduction of long chain branching by the use of a supported bis-cyclopentadienyl transition metal compound having an alkylene or silyl bridge used in conjunction with a methylalumoxane cocatalyst. These catalysts are supported. The batch reactions are with a scavenger (see page 5 line 28). Example 10 of this patent publication discloses an Melt Index (MI) of 0.3 g per 10 minutes determined at 190 C under a 2.16 kg load; there is no indication of the molecular weight distribution, the Compositional Distribution is not given, the density is 20 0.916 g/ml, the Haze is 11 %, there is no indication of the ratio of MI's determined under different loads, the Dart Impact Strength is 210 g/mil and there is no indication of the polymer stiffness as expressed by the modulus. On the basis of the correlation between density and secant modulus given in the Encyclopedia of Polymer Science and Engineering, by Mark, Bikales, Overberger, and Menges, Vol. 6, second ed., p. 447 (1986), the secant modulus for this material is estimated to be about 30,000 to 32,000 psi (205 to 220 N/mm²).

EP-A-659773 (Union Carbide) used bridged catalysts with examples running the reactor in a continuous mode in the polymerization of polyethylene. The specification teaches the use of support (see page 6

line 30) but the examples do not use a support so that the alumoxane is in a solvent when injected. This may favor fouling and furthermore the alumoxane will contain a significant amount of unreacted trimethyl aluminum (TMA) which may act as a scavenger and lead to an apparent 5 increase in vinyl unsaturation. Melt processing is further influenced by the use of more than one metallocene component which can broaden the molecular weight distribution by the production of more than one distinct polymer component. This is done allegedly to provide control over the degree of long chain branching (LCB) as indicated by the degree of LCB determined by GPC and viscosity data. The melt flow ratio (MFR) is also used to characterize the polymer. The MFR is the ratio of melt index (MI) at different loads and reflects LCB and higher Mw/Mn. Increasing MFR values may be due to higher Mw/Mn caused by the use of more than one metallocene. The examples indicate that the bridged species is most instrumental in raising the level of LCB. However, Example 5 shows the use of the bridged metallocene alone produces a polymer having a very low molecular weight, suggesting that the low molecular weight polymer species are a major contributor to higher MFR values. EP-A-659773 thus fails to teach how a low melt index material may be produced which has 20 the improved rheology as expressed in MFR resulting from the presence of LCB. EP-A-659773 does not disclose the CDBI, haze and DIS values which help determine the commercial quality of the polymer produced.

EP-A-743327 describes the preparation of an ethylene polymer having a high polydispersity index (which equals Mw/Mn) which requires a lower head pressure in extrusion. The improved rheological properties are expressed in terms of RSI (Relaxation Spectrum Index) which is said to be sensitive to molecular weight distribution, molecular weight and long chain branching. The polymerization process details are scant. EP-A-743327 includes as catalyst similar metallocenes to those listed in EP-A-659773.

WO 98/44011 PCT/US98/06081

-5-

EP-A-729978 characterizes an ethylene polymer using flow activation energy. The polymer is made using bridged bis cyclopentadienyl catalyst components, with one cyclopentadienyl ring system being a fluorenyl polynuclear ligand structure. The higher activation energy may be the result of higher levels of long chain branching.

Many different process or catalyst options are introduced in the above processes to achieve the desired effect in the melt processing of the resulting polymers. However it is suggested that these processes all suffer from drawbacks which mitigate against commercial implementation in that the catalyst may have low productivity, be prone to fouling in the longer runs used for large scale reactors and/or produce low molecular weight materials. In addition the prior proposals may lead to an undue sacrifice of physical properties such as loss of clarity, increase in extractability which is detrimental in food contact applications, or loss of film toughness properties such as DIS (dart impact strength).

It is amongst the aims of the invention to provide a relatively simple process for providing commercially desirable polymer from commercial scale plants which has advantageous melt flow properties and balance of strength and stiffness.

The polymer can be produced in prolonged production runs under conditions not likely to lead to fouling.

SUMMARY OF THE INVENTION

The invention provides a polymer of an ethylene and at least one alpha olefin having at least 5 carbon atoms obtainable by a continuous gas phase polymerization using supported catalyst of an activated molecularly discrete catalyst, such as a metallocene, in the substantial absence of an

WO 98/44011 PCT/US98/06081

-6-

aluminum alkyl based scavenger (e.g., triethylaluminum (TEAL), trimethylaluminum (TMAL), tri-isobutylaluminum (TIBAL), tri-n-hexylaluminum (TNHAL) and the like), which polymer has a Melt Index (MI) as herein defined of from 0.1 to 15; a Compositional Distribution Breadth Index (CDBI) as defined herein of at least 70%, a density of from 0.910 to 0.930 g/ml; a Haze value as herein defined of less than 20; a Melt Index ratio (MIR) as herein defined of from 35 to 80; an averaged Modulus (M) as herein defined of from 20 000 to 60 000 psi (pounds per square inch) (13790 to 41369 N/cm²) and a relation between M and the Dart Impact Strength in g/mil (DIS) complying with the formula:

$$DIS \ge 0.8 \times [100 + e^{(11.71 - 0.000268 \times M + 2.183 \times 10^{-9} \times M^2})],$$

where "e" represents 2.1783 the base Napierian logarithm, M is the averaged Modulus in psi, and DIS is the 26 inch (66cm) dart impact strength in g/mil.

15

While many prior art documents describe processes and polymers using the same monomers and similar processes, none describe polymers combining [A] good shear thinning and therefore relatively favorable extrusion and other melt processing properties with [B] a high stiffness and [C] high impact strength. Up to now, these features appeared to be difficult to combine in LLDPE (linear low density polyethylene) materials produced in a continuous gas phase process. The invention provides a surprising combination of properties for the polymer which can be prepared reproducibly.

25

30

In comparison to LDPE (low density polyethylene) made in the high pressure process having a comparable density and MI, the polyethylenes of the invention have a favorable DIS-Modulus balance, e.g., a dart impact strength (DIS) in g/mil that is equal to or greater than that predicted by the formula:

DIS
$$\ge$$
0.8 x [100+e^{(11.71-0.000268xM+2.183x10-9xM²)],}

where "e" is the base Napierian logarithm, M is the averaged Modulus in psi, and DIS is the dart impact strength in g/mil.

In comparison with LLDPE made by a gas phase process using conventional Ziegler Natta supported catalysts, the polyethylenes of the invention have improved shear thinning. These conventionally produced LLDPE's will have a relatively low CDBI and a poor DIS-Modulus balance, e.g., a dart impact strength in g/mil that is less than that predicted by the above formula.

10

In comparison to the EXCEED™ materials (made by Exxon Chemical) produced in gas phase processes using metallocene based supported catalysts, the polyethylenes of the invention have a better shear thinning behavior and comparable other properties. The MIR will be from 16 to 18 for such EXCEED materials.

In a preferred form of the invention, the polyethylenes of the invention are derived from ethylene and up to 15 weight percent of 1-hexene. Preferably the relation between the Modulus and the Dart Impact Strength complies with the formula:

$$\label{eq:discrete_discrete_discrete_bound} \text{DIS} \leq 2.0 \; \text{x} \; [\; 100 + e^{\left(11.71 - 0.000268 \text{xM} + 2.183 \text{x} 10^{-9} \text{x} \text{M}^2\right)}],$$

where "e" is the base Napierian logarithm, M is the averaged Modulus in psi, and DIS is the dart impact strength in g/mil.

Advantageously the polymer may have either one or combination of the following features: the density is from 0.915 to 0.927 g/ml, the MI is from 0.3 to 10 and CDBI is at least 75 %. Most preferred is a DIS is from 120 to 1000 g/mil, especially less than 800 and more than 150 g/mil. Preferably the Mw/Mn by GPC is from 2.5 to 5.5

WO 98/44011

-8-

As to the process conditions the overall conditions described in WO 96/08520 (Exxon Chemical Patents Inc) can be adopted. Inventors believe that a combination of particular process conditions helps to make the polyethylene of the invention. In particular, it is thought desirable to use a catalyst system in which the metallocene has a pair of bridged cyclopentadienyl groups, preferably with the bridge consisting of a single carbon, germanium or silicon atom so as to provide an open site on the catalytically active cation. The activator may be methyl alumoxane as described in EP-129368 or a noncoordinated anion as described in EP-277004. It also thought desirable that there should be substantially no scavengers which may interfere with the reaction between the vinyl end unsaturation of polymers formed and the open active site on the cation. By the statement "substantially no scavengers", it is meant that there should be less than 100 ppm by weight of such scavengers present in the feed gas, or preferably no intentionally added scavenger, e.g., an alkylaluminum or Lewis acidic scavenger, other than that which may be present on the support.

The conditions optimal for the production of the polyethylene of the invention also require steady state polymerization conditions which are not likely to be provided by batch reactions in which the amounts of catalyst poisons can vary and where the concentration of the comonomer may vary in the production of the batch.

25

Overall continuous gas phase process for the polymerization of a polyethylene may thus comprise:

continuously circulating a feed gas stream containing monomer and inerts to thereby fluidize and agitate a bed of polymer particles, adding metallocene catalyst to the bed and removing polymer particles in which: a) the catalyst comprises at least one bridged bis cyclopentadienyl transition metal and an alumoxane activator on a common or separate porous support;

 b) the feed gas is substantially devoid of a Lewis acidic scavenger and wherein any Lewis acidic scavenger is preferably present in an amount

less than 100 wt. ppm of the feed gas;

- the temperature in the bed is no more than 20°C less than the polymer melting temperature as determined by DSC, at a ethylene partial pressure in excess of 60 pounds per square inch absolute (414 kPaa), and
- d) the removed polymer particles have an ash content of transition metal of less than 500wt.ppm, the MI is less than 10, the MIR is at least 35 with the polymer having substantially no detectable chain end unsaturation as determined by HNMR
- By the statement that the polymer has substantially no end chain unsaturation, it is meant that the polymer has vinyl unsaturation of less than 0.1 vinyl groups per 1000 carbon atoms in the polymer, e.g., less than 0.05 vinyl groups per 1000 carbons, e.g., 0.01 vinyl groups per 1000 carbons or less.
- The process aims to provide the polyethylene of the invention throughout the use of a single catalyst and the process does not depend on the interaction of bridged and unbridged species. Preferably the catalyst is substantially devoid of a metallocene having a pair of pi bonded ligands (e.g., cyclopentadienyl compounds) which are not connected through a covalent bridge, in other words, no such metallocene is intentionally added to the catalyst, or preferably, no such metallocene can be identified

in such catalyst, and the process uses substantially a single metallocene species comprising a pair of pi bonded ligands at least one of which has a structure with at least two cyclic fused rings (e.g., indenyl rings). Best results may be obtained by using a substantially single metallocene species comprising a monoatom silicon bridge connecting two polynuclear ligands pi bonded to the transition metal atom.

The catalyst is preferably supported on silica with the catalyst homogeneously distributed in the silica pores. Preferably, fairly small amounts of methyl alumoxane should be used, such as amounts giving an Al to transition metal ratio of from 400 to 30, and especially of from 200 to 50.

In order to obtain a desired melt index ratio, the molar ratio of ethylene and comonomer can be varied, as can concentration of the comonomer. Control of the temperature can help control the MI. Overall monomer partial pressures may be used which correspond to conventional practice for gas phase polymerization of LLDPE.

The parameters used in the claims and the examples are defined as follows

Melt Index:

ASTM D-1238- Condition E

25 Melt Index ratio:

this is ratio of l21 over l2 as determined

by ASTM D-1238.

Mw, Mn and Mw/Mn: determined by GPC using a DRI (differential refraction index) detector.

Gel permeation chromatography (GPC) is performed on a Waters 150C GPC instrument with DRI detectors.

GPC Columns are calibrated by running a series of narrow polystyrene standards. Molecular weights of polymers other than polystyrenes are conventionally calculated by using Mark Houwink coefficients for the polymer in question.

CDBI is determined as set out in column 7 and 8 of WO9303093.

10

SCB (short chain branching): This was determined by HNMR (hydrogen nuclear magnetic resonance) with data collected at 500 Mhz. Spectra were referenced by setting the polymer backbone signal to 1.347 ppm. Methyl group contents in ethylene 1-olefin copolymers were calculated from the HNMR spectrum using the following formula:

Methyl Groups/1000 Carbons = $(I_{CH3}*0.33*1000)/(I_{0.5-}_{2.1ppm}*0.5)$

20

30

where l_{CH3} is the normalized methyl signal area in the region between 0.88 and 1.05 ppm and $l_{0.5\text{-}2.1\text{ppm}}$ the area between 0.50 and 2.10 ppm.

The amount of methyl groups will correspond to the number of short chain branches in the polymer assuming that the short chain branches contain 1 methyl (-CH₃) group and that all methyl groups are a result of short chain branching. The same NMR method can be used to determine vinyl end unsaturation.

Density:

ASTM D-1505

WO 98/44011

Haze %:

ASTM D-1003-95

Dart Impact Strength, 26 inch (66cm):

ASTM D1709-91

5

1% secant Modulus:

ASTM D-882-91

The "averaged Modulus" is the sum of the 1% secant Modulus in the machine direction and in the transverse direction divided by two.

10

Elmendorf tear strength

ASTM D1922-94

Granular Bulk Density: The granular polymer particles are poured via a 7/8 " diameter funnel into a fixed volume cylinder of 400 ml. The bulk density is measured as the weight of resin divided by 400 ml to give a value in g/ml.

Particle Size: The particle size is measured by determining the weight of material collected on a series of U.S. Standard sieves and determining the weight average particle size in micrometers based on the sieve series used.

Extractability: determined according to FDA regulations 21CFR 177.1520 (d) (3) (ii).

25 **EXAMPLES**

Two runs are illustrated in detail below.

Catalyst preparation

Run 1 Supported Catalyst Preparation:

30 A solution of 1300 ml of 30 wt% alumoxane (MAO) in toluene as determined by reference to the total Al content which may include

unhydrolyzed TMA, was charged to a two gallon (7.57 Liter), jacketed glass-walled reactor, equipped with a helical ribbon blender and an augertype shaft. 2080 ml of toluene was added and stirred. A suspension of dimethylsilyl-bis-(tetrahydroindenyl) zirconium dichloride 31.5 (Me₂Si(H₄Ind)₂ZrCl₂) in 320 ml of toluene purchased from Albemarle Labs, was cannulated to the reactor. An additional bottle of dry toluene (250 ml) was used to rinse solid metallocene crystals into the reactor by cannula under nitrogen pressure. A color change from colorless to vellow/orange was noted upon addition of the metallocene to the MAO solution. The mixture was allowed to stir at 69°F (20.6°C) for one hour, before being transferred to a four-liter Erlenmeyer flask under nitrogen. Silica (1040 g, Davison MS 948, 1.65 ml/g pore volume was charged to the reactor. Half of the solution from the 4 liter Erlenmeyer flask was then transferred back to the 2 gallon (7.57 liter) stirred glass reactor. The reaction temperature rose from 70°F (21.1 °C) to 100°F (37.8°C) in a five minute exotherm. The balance of the solution in the 4 liter Erlenmeyer was subsequently added back to the glass reactor, and stirred twenty minutes. Then, toluene was added (273 ml, 238 g) to dilute the active catalyst slurry, and stirred an additional twenty-five minutes. Antistat AS-990, a surface modifier made from ethoxylated stearylamine sold by Witco Chemical Corp. (7g in 73 ml toluene) was cannulated to the reactor and the slurry mixed for thirty minutes. Removal of solvent commenced by reducing pressure to less than 18 inches of mercury (457 mmHg) while feeding a small stream of nitrogen into the bottom of the reactor and raising the temperature from 74°F (23.3 °C) to 142°F (61.1 °C) over a period of one hour. Then five additional hours of drying at 142°F (61.1 °C) to 152°F (66.7 °C) and vacuum which ranged from 5 inches to 22 inches Hg (127 to 559 mmHg) were used to dry the support and yield 1709.0 g of free-flowing active supported catalyst material. Head space gas chromatograph (HSGC) measurements showed 13,000 weight parts

30

per million (1.3 wt %) of residual toluene. A second drying step under stronger vacuum conditions, resulted in HSGC analysis measurement of residual toluene at 0.18%. Elemental analysis showed 0.40% Zr, 10.75% Al, 30.89% Si, 0.27% Cl, 9.26% C, 2.05% H (all percentages shown herein are weight percent).

Run 2 Supported Catalyst Preparation:

A solution of 1125 ml of 30 wt% alumoxane (MAO) in toluene as determined by reference to the total Al content which may include 10 unhydrolyzed TMA was charged to a two gallon (7.57 liter), jacketed glass-walled reactor, equipped with a helical ribbon blender and an augertype shaft. 1800 ml of toluene was added and stirred. A suspension of zirconium dichloride dimethylsilyl-bis-(tetrahydroindenyl) (Me2Si(H4Ind)2ZrCl2) in 320 ml of toluene purchased from Albemarle 15 Labs, was cannulated into the reactor. An additional 150 ml of toluene was used to rinse solid metallocene crystals into the reactor by cannula under nitrogen pressure. A color change from colorless to yellow/orange was noted upon addition of the metallocene to the MAO solution. The mixture was allowed to stir at 69°F (20.6°C) for one hour, before being transferred to a four-liter Erlenmeyer flask under nitrogen. Silica (899 g, Davison MS 948, 1.65 ml /g Pore Volume, V.) was charged to the reactor. Half of the solution from the 4 L Erlenmeyer flask was then transferred back to the 2 gallon (7.57 liter) stirred glass reactor. The reaction temperature rose from 70°F (21.1 °C) to 100°F (37.8°C) in a five minute 25 exotherm. The balance of the solution in the 4 liter Erlenmeyer was subsequently added back to the glass reactor, and stirred twenty minutes. Then, toluene was added (273 ml, 238 g) to dilute the active catalyst slurry, and stirred an additional twenty-five minutes. Antistat AS-990, a surface modifier sold by Witco (6 g in 30 ml toluene), was cannulated to the reactor and the slurry mixed for thirty minutes. Removal of solvent commenced by reducing pressure to less than 18 inches of mercury (457

mmHg) while feeding a small stream of nitrogen into the bottom of the reactor and raising the temperature from 74°F (23.3 °C) to 142°F (61.1 °C) over a period of one hour. Then nine and a half additional hours of drying at 142°F (61.1 °C) to 152°F (66.7 °C) at a vacuum which ranged from 5 inches to 22 inches Hg (127 to 559 mmHg) were used to dry the support and yield 1291.4 g of free-flowing active supported catalyst material.

Fluid-Bed Polymerization:

The polymerization was conducted in a continuous gas phase fluidized bed reactor having a 16.5 inch (41.9cm) diameter with a bed height of approximately 12 feet (3.6m). The fluidized bed is made up of polymer granules. The gaseous feed streams of ethylene and hydrogen together with liquid comonomer were mixed together in a mixing tee arrangement and introduced below the reactor bed into the recycle gas line. The individual flow rates of ethylene, hydrogen and comonomer were controlled to maintain fixed composition targets. The ethylene concentration was controlled to maintain a constant ethylene partial pressure. The hydrogen was controlled to maintain a constant hydrogen to ethylene mole ratio. The concentration of all the gases were measured by an on-line gas chromatograph to ensure relatively constant composition in the recycle gas stream.

The solid catalyst was injected directly into the fluidized bed using purified nitrogen as a carrier. Its rate of injection was adjusted to maintain a constant production rate of the polymer. The reacting bed of growing polymer particles is maintained in a fluidized state by the continuous flow of the make up feed and recycle gas through the reaction zone. A superficial gas velocity of 1-3 ft/sec (0.3 to 0.9 m/sec) was used to achieve this. The reactor was operated at a total pressure of 300 psig (2068 kPa gauge). To maintain a constant reactor temperature, the

temperature of the recycle gas is continuously adjusted up or down to accommodate any changes in the rate of heat generation due to the polymerization.

The fluidized bed was maintained at a constant height by withdrawing a portion of the bed at a rate equal to the rate of formation of particulate product. The product is removed semi-continuously via a series of valves into a fixed volume chamber, which is simultaneously vented back to the reactor. This allows for highly efficient removal of the product, while at the same time recycling a large portion of the unreacted gases back to the reactor. This product is purged to remove entrained hydrocarbons and treated with a small stream of humidified nitrogen to deactivate any trace quantities of residual catalyst and cocatalyst.

TABLE 1

15

Polymerization Run	Run 1	Run 2
Condition		
Zr (wt %)	0.43	0.50
Al (wt %)	11.6	11.4
Al/Zr (mole/mole)	91.2	77.1
Temperature (°C)	79.4	85
Pressure (bar)	21.7	21.7
Ethylene (mole %)	25.0	49.9
Hydrogen (mole ppm)	275	445
Hexene (mole %)	0.23	0.32
Bed Weight (Kg PE)	113	121
Production Rate (Kg	27.6	35.5
PE/Hr)		
Catalyst Productivity	1690	2287
(Kg PE/Kg catalyst)		

Bulk Density (g/ml)	0.448	0.450
Average Particle Size	920	803
(micronmeters)		
Ash (ppm)	507	386

The parameters were determined as set out previously; the Zr, Al wt percent and ash levels were by elemental analysis.

No aluminum alkyl compounds were added to the reactor as scavenger. The runs were continued for around 3 days.

The polymers resulting were subjected to additional tests, where appropriate after first forming the polymer into film.

Polymer characterization

10

TABLE 2A

	Run 1	Run 2
Density	0.9190	0.9257
MI	1.10	0.62
MIR	46.0	57.6
DRI Detector		
Mw	92,200	104700
Mn	18,300	17900
Mz	208,400	287500
Mw/Mn	5.04	5.85

TABLE 2A continued

	Run 1	Run 2	••
DSC 2nd Melt-See note 1		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	••
1st melting peak. (C)	108.6	122.6	
2nd. melting peak. (C)	119.3	117.3	
CDBI	86	83.10	
SCB (/1000 C)	15.4	10.6	
wt% C6	9.3	6.4	
mole% C6	3.3	2.2	

Note1: the sample had been molten and been allowed to cool once previously

Table 2B film characterization

	Run 1	Run 2
Blow up ratio	2.5	2.5
Gauge in mil (1 mil=25.4micronmeter)	2.1	2.0
1% Secant Modulus, psi (N/cm²) MD(machine direction)	29420 (20284)	45070 (31075)
TD (transverse direction), psi (N/cm²)	31230 (21532)	47420 (32695)
MD+TD average, psi (N/cm²)	30325 (20908)	46245 (31885)
Elmendorf tear strength (g/mil)	•	
MD	207	134
TD	430	477
26 inch (66cm) Dart Impact Strength (DIS) in g/mil	410	156

Calculated DIS as a function of Modulus as per the formula	386	154
HAZE (%)	10.2	9.9
Extractability	1.0	not
		available

A larger number of further tests were performed with different samples made according to the invention in a similar manner was made and the results are shown in the Drawing 1.The function in the claim 1 is shown as a solid line.

Table 3 shows some exemplary values.

TABLE 3

Average	Averaged 1% Secant Dart Impact Strength (26 inch) (66cm)		nch) (66cm)
Modulus, M as Measured		•	
psi	N/cm²	Calculated from Formula	Measured
		DIS≥0.8 x [100+e ^{(11.71} -	g/mil
		0.000268xM+2.183x10 ⁻⁹ xM ²)]	
		g/mil	
25,575	17,633	508	611
28,580	19,705	353	456
28,990	19,987	337	553
29,145	20,094	332	451
30,325	20,908	294	410
31,450	21,684	264	284
31,610	21,794	260	257
32,000	22,063	. 251	349

32,140	22,159	248	223
32,140	22,109		
33,780	23,290	217	251
34,160	23,552	211	262
35,170	24,248	196	223
35,970	24,800	186	261
37,870	26,110	167	251
39,325	27,113	155	197
39,390	27,158	154	193
43,675	30,112	131	167
46,245	31,884	123	156
47,730	32,908	119	147
49,460	34,101	115	143

The claims therefore cover the combination of DIS and averaged Modulus in the area of Fig. 1 above the solid line.

Using the indications and guidance provided in the specification concerning catalyst selection, catalyst support and gas phase process operation it is possible to produce ethylene polymers as specified in the claims which are simultaneously optically clear; relatively easy to make and to process and have a high strength as measured by the Dart Impact Strength.

The films can be used for heavy duty bags, shrink film, agricultural film, particularly which are down-gauged such as garbage and shopping bags with a thickness of from 0.5 to 7 mil. The films can be produced by blow extrusion, cast extrusion, co-extrusion and be incorporated also in laminated structures.

Claims

20

We Claim:

1. A polymer of ethylene and at least one alpha olefin having at least 5 carbon atoms, which polymer has a Melt Index (MI) as herein defined of from 0.1 to 15; a Compositional Distribution Breadth Index (CDBI) as defined herein of at least 70%, a density of from 0.910 to 0.930 g/ml; a Haze value as herein defined of less than 20 %; a Melt Index ratio (MIR) as herein defined of from 35 to 80; an averaged Modulus (M) as herein defined of from 20 000 to 60 000 psi (pounds per square inch) (13790 to 41369 N/cm²) and a relation between M and the Dart Impact Strength in g/mil (DIS) complying with the formula:

$$DIS \ge 0.8 \times [100 + e^{(11.71 - 0.000268 \times M + 2.183 \times 10^{-9} \times M^2})].$$

2. The polymer according to claim 1 in which the relation between the averaged Modulus and the Dart Impact Strength (DIS) complies with the formula:

DIS
$$\leq 2.0 \times [100 + e^{(11.71 - 0.000268 \times M + 2.183 \times 10^{-9} \times M^2]}$$
.

- 3. The polymer according any of the preceding claims in which the density is from 0.915 to 0.927 g/ml, the MI is from 0.3 to 10 and the CDBI is at least 75 %.
- 4. The polymer according any of the preceding claims in which the DIS is from 120 to 1000 g/mil.
- 5. The polymer according any of the preceding claims in which the levels of extractables as defined herein is less than 2.6 wt %.
 - A film comprising at least one layer of the polymer according to any of the preceding claims.

- 7 The film according to claim 6 in which the layer has a thickness of from 0.5 to 7 mil and is preferably a mono-layer.
- 5 8 A continuous gas phase process for the polymerization of a polyethylene according to any of the preceding claims 1 to 4 comprising:

continuously circulating a feed gas stream containing monomer and inerts to thereby fluidize and agitate a bed of polymer particles, adding metallocene catalyst to the bed and removing polymer particles in which:

the catalyst comprises at least one bridged bis cyclopentadienyl transition metal and an alumoxane activator on a common or separate porous support;

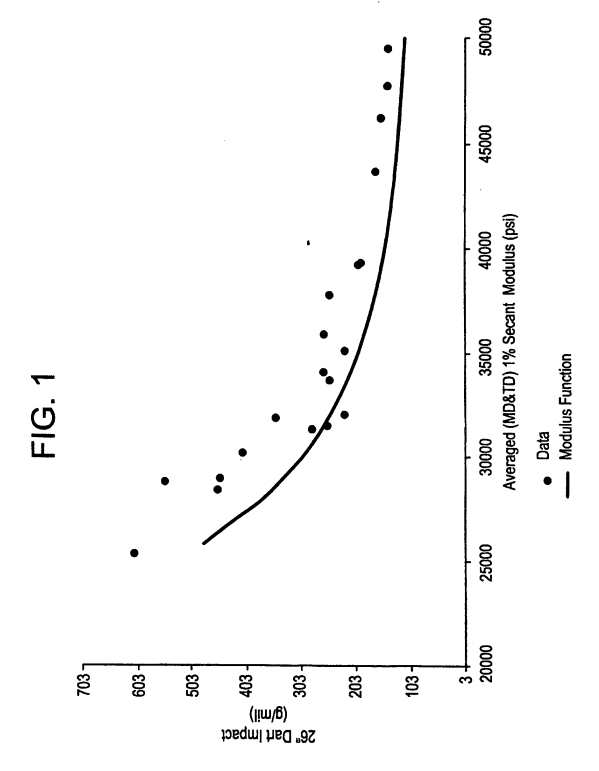
the feed gas is substantially devoid of a Lewis acidic scavenger and wherein the Lewis acidic scavenger is preferably present in an amount less than 100 ppm of the feed gas;

the temperature in the bed is no more than 20°C less than the polymer melting temperature as determined by DSC, at an ethylene partial pressure in excess of 75 pounds per square inch absolute (517 kPaa) and

the removed polymer particles have an ash content of transition metal of less than 500wt.ppm, the MI is less than 10, the MIR is at least 40 with the polymer having substantially no detectable chain end unsaturation.

9. The process according to claim 8 in which the catalyst is substantially devoid of a metallocene having a pair of pi bonded ligands which are not connected through a covalent bridge.

- 10. The process according to claim 8 or 9 wherein there is used a substantially single metallocene species comprising a pair of pi bonded ligands at least one of which has a structure with at least two cyclic fused 5 rings.
- 11. The process according to any of claims 8 to 10 wherein there is used a substantially single metallocene species comprising a mono-atom silicon bridge connecting two polynuclear ligands pi-bonded to the transition metal atom.



Inte. onal Application No PCT/US 98/06081

A. CLASSI IPC 6	COSF210/16 COSF2/34 COSF4	1/602 C08J5/18	
According to	o International Patent Classification(IPC) or to both national cla	ssification and IPC	
	SEARCHED		
Minimum do IPC 6	ocumentation searched (classification system followed by class COSJ COSF	ification symbols)	
Documenta	tion searched other than minimum documentation to the extent	that such documents are included in the fields sec	arched
Electronic d	data base consulted during the international search (name of da	ata base and. where practical, search terms used	
С. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the	he relevant passages	Relevant to claim No.
X	WO 94 14855 A (MOBIL OIL CORP see examples 1-3; table II see claim 1	•	1,3,5,6
Х	see page 1, line 12 - line 13	; claim 4	4
X	EP 0 676 421 A (BP CHEM INT L October 1995 cited in the application see example 17 : page 4 line ! line 24-45; table 1 page 10		8-11
х	EP 0 685 496 A (MITSUI PETROCI 6 December 1995 see example 3	HEMICAL IND)	8-10
		-/	
X Furt	lher documents are listed in the continuation of box C.	χ Patent family members are listed	in annex.
3 Special ca	ategories of cited documents :	"T" later document published after the inte	rnational filing date
consider consider consider the consider	ent which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another on or other special reason (as specified) sent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but than the priority date claimed	or priority date and not in conflict with cited to understand the principle or th invention "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the dc "Y" document of particular relevance; the cannot be considered to involve an in document is combined with one or ments, such combination being obvious the art. "&" document member of the same patent	the application but servi underlying the claimed invention to considered to coursent is taken alone claimed invention aventive step when the ore other such docurus to a person skilled
	actual completion of theinternational search	Date of mailing of the international sea	arch report
	Page 1998 Mailing address of the ISA	09/07/1998 Authorized officer	
	European Patent Office. P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Gamb, V	

1

Int. donal Application No PCT/US 98/06081

	PCT/US 98/06081
ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
WO 97 29138 A (EXXON CHEMICAL PATENTS INC) 14 August 1997 see example 6 see claims 10,11,15 see page 11, line 28 - line 32 see page 12, line 9 - line 17	8-11
WO 98 02483 A (SENTINEL PRODUCTS CORP) 22 January 1998 see page 32, line 24 - page 33, line 15 : éxact 3012 and 3011	1,3
EP 0 374 783 A (VISKASE CORP) 27 June 1990 see example 5 table 2 page 15 and page 14 line 8-9	1,3,6
·	
	WO 97 29138 A (EXXON CHEMICAL PATENTS INC) 14 August 1997 see example 6 see claims 10,11,15 see page 11, line 28 - line 32 see page 12, line 9 - line 17 WO 98 02483 A (SENTINEL PRODUCTS CORP) 22 January 1998 see page 32, line 24 - page 33, line 15: éxact 3012 and 3011 EP 0 374 783 A (VISKASE CORP) 27 June 1990 see example 5 table 2 page 15 and page 14

1

information on patent family members

Int. ional Application No PCT/US 98/06081

Patent document			Publication		atent family	Publication
cited in	n search report		date	member(s)		date
WO 9	9414855	Α		ŞG	46293 A	20-02-1998
				US	5332706 A	26-07-1994
				US	5420220 A	30-05-1995
				AU	5213298 A	23-04-1998
				ΑU	682842 B	23-10-1997
				AU	5749294 A	19-07-1994
				AU	673048 B	24-10-1996
				AU	5871594 A	19-07-1994
				AU	672905 B	17-10-1996
				AU	6551394 A	11-10-1994
				CA	2152623 A	07-07-1994
				CA	2152937 A	07-07-1994
				CA	2156528 A	29-09-1994
				EP	0675907 A	11-10-1995
				EP	0675906 A	11-10-1995
				EP	0690878 A	10-01-1996
				JP	8505172 T	04-06-1996
				JP	8505174 T	04-06-1996
				JP	8508304 T	03-09-1996
				WO WO	9414856 A	07-07-1994
				US	9421691 A	29-09-1994
				US	5608019 A 5473028 A	04-03-1997 05-12-1995
				US	5602067 A	11-02-1997
			~~~~~~~~~~~~~			11-02-1997
EP (	676421	Α	11-10-1995	CA	2146205 A	08-10-1995
		<b></b>		JP	8048711 A	20-02-1996
EP C	685496	Α	06-12-1995	JP	2173014 A	04-07-1990
				JP	2685261 B	03-12-1997
				JP	2173016 A	04-07-1990
				JP	2685262 B	03-12-1997
				JP	2173015 A	04-07-1990
				JP	2685263 B	03-12-1997
				EP	0685498 A	06-12-1995
				AT	166890 T	15-06-1998
				EP	0495099 A	22-07-1992
				WO	9007526 A	12-07-1990
				US	5525689 A	11-06-1996
				US	5714426 A	03-02-1998
			•			

Information on patent family members

Int. Jonal Application No PCT/US 98/06081

Patent document cited in search report			Publication date	Patent family member(s)		Publication date
EP (	0685496	A		US US EP US CA JP JP	5218071 A 5336746 A 0769505 A 5639842 A 2008315 A,C 2276807 A 2571280 B	08-06-1993 09-08-1994 23-04-1997 17-06-1997 24-07-1990 13-11-1990 16-01-1997
WO 9	9729138	Α	14-08-1997	NONE		
WO S	9802483	A	22-01-1998	AU	3600797 A	09-02-1998
EP (	0374783	A	27-06-1990	AU AU CA JP JP US	634991 B 4685289 A 2003882 A,C 2229835 A 2573527 B 5434010 A	11-03-1993 21-06-1990 19-06-1990 12-09-1990 22-01-1997 18-07-1995